Visual-vestibular Interaction: Basic Science to Clinical Relevance

Summary
The visual and vestibular systems together mediate the reflex and perceptual functions required for efficient postural balance and spatial orientation in the light or the dark. Following an acute unilateral vestibular insult, there is a left-right imbalance in the vestibular input leading to erroneous brainstem vestibular signaling. This asymmetrical vestibular signaling manifests in several ways, e.g. activation of reflex eye movements (vestibular nystagmus) and sensations of vertigo. The net result is a functional impairment in the capacity to locomote and spatially orientate; a high risk situation in a natural environment. The central nervous system can however promote a rapid functional recovery by shifting the brain’s relative reliance from vestibular toward visual cues for these functions, i.e. the brain becomes more ‘visually dependent’ for the sensory monitoring of locomotion and spatial orientation. Given time and the appropriate conditions (e.g. adequate physical activity, avoidance of vestibular sedatives), the fidelity of the brainstem vestibular signal is restored as the brain process involved in ‘vestibular compensation’ removes the left-right imbalance in the vestibular brainstem signal. In patients who make a full symptomatic recovery consequent upon adequate ‘vestibular compensation’, the reliance on visual signals for locomotion and navigation reduces toward normal levels in tandem with the process of brainstem vestibular compensation. In contrast, some chronically symptomatic patients show a maladaptive persistence of this heightened visual dependency and this is manifest in visually-induced dizziness (“visual vertigo”). Visually-induced dizziness is a major problem in the clinic, with symptoms occurring in visually-busy environments such as shopping malls or supermarkets. We discuss the physiological mechanisms underlying visual-vestibular interaction, how this interaction may be disturbed leading to visually-induced dizziness and finally how understanding the physiological mechanism helps in the development of therapy for these patients.

Introduction
The vestibular system, which provides a signal of head motion to the brain, mediates functions of gaze and postural stabilisation via vestibulo-ocular reflex (VOR) and vestibulospinal reflexes. The vestibular system is also key in generating sensations of self-motion and spatial orientation required for navigation in the environment. The vestibular system influences these reflex and perceptual functions in partnership with other sensory systems, particularly vision. For example, vision calibrates the accuracy of the VOR and, via optic flow and motion parallax generated during self-motion, contributes to our sense of self-motion (or stasis). Occasionally, visual-vestibular interaction can mislead, e.g. the compelling but false sensation of self-motion experienced when looking out of the window of a stationary train as an adjacent train moves past us (‘the ‘train illusion’, Figure 1A). This illusion demonstrates the difficulty the brain has in trying to resolve the complex and ambiguous role of the visual system in signalling both self- and object (environmental) motion.

The occasional failure of the normal brain to accurately estimate measures of self-motion is key to understanding how many patients’ symptoms relate to an abnormal visual-vestibular interaction. Indeed, patients with vestibular disorders commonly report a modulation of their dizziness by visual stimuli. In acute vertigo, when typically there are abnormal signs such as a vestibular nystagmus, patients often close their eyes to avoid the distressing illusion of seeing the world spinning. In contrast, in chronic dizziness where there are usually no abnormal signs (an apparent uncoupling of symptoms from signs), patients complain of dizziness in the face of relatively trivial motion in the environment (e.g. crowds in a shopping mall). Since such visual stimuli are ubiquitous in the modern world, visually-induced dizziness (so-called ‘visual vertigo’) may be crippling for patients’ social, occupational and mental well-being. In this overview we explore the basic mechanisms underlying the intimate relationship between visual motion and dizziness and the relevance of this visual-vestibular interaction for patients’ symptoms and their management.

Role of the vestibular system in health
As we navigate through the environment, our visual system is faced with two challenges: (1) the maintenance of a stable and clear image of the world during head movements; (2) the accurate ascribing of visual motion as being due to either self-motion or environmental motion – put simply the brain asks the question: am I moving? or is the object/world moving? To overcome these problems the central nervous system combines visual and vestibular inputs.

Maintaining a clear and stable vision is enabled by a natural ‘steady-cam’ mechanism called the vestibulo-ocular reflex (VOR). The VOR involves a 3-neurone brainstem reflex that begins with the detection of head acceleration by the peripheral
labyrinth. This head motion signal is conveyed by primary vestibular afferents to the vestibular nuclei neurones in the brainstem which in turn project to ocular motor neurones. The VOR thus keeps the eyes steady and ‘locked on’ to the visual target of interest despite head motion. This mechanism thus maintains visual acuity and a stable visual world by reducing slippage of the visual image across the retina. This ‘retinal slip’, when it does occur, may provoke the unpleasant sensation of oscillopsia. In general the degree of oscillopsia is coupled to the amount of retinal slip, particularly in the acute state. Retinal slip and oscillopsia symptoms are not however inevitably linked but can be uncoupled in the chronic adapted state. The capacity for the brain to render a physical retinal slip unnoticeable is an important concept since it leads to the finding that ocular motor (reflex) parameters of vestibular function (i.e. VOR) relate poorly to perceptual aspects of vestibular function (i.e. dizziness) in the chronic state. Indeed a relatively common but extreme example of such perceptuo-reflex uncoupling is that seen in idiopathic congenital nystagmus where a vigorous nystagmus is not associated with symptoms.

The visuo-vestibular interaction

One mechanism proposed to solve the motion ambiguity problem is that of a reciprocal visual-vestibular inhibition (Figure 2). Specifically if the vestibular system signals ‘no motion’, then this impedes a visual motion signal from indicating self-motion. Conversely, when there is no vestibular signal, visual input can provoke a sensation of self-motion but only if the visual stimulus occupies a sufficiently large visual area. Cognitive influences are also important since illusory self-motion is more likely to occur if there is a high probability of self-motion, e.g. sitting on a train is a situation where motion is likely, whereas sitting on the sofa has a low probability of motion. The psychophysical evidence for a visuo-vestibular reciprocal inhibition is supported by behavioural data showing that during self-motion, the threshold for visually detecting object-motion is elevated. Similarly visual-vestibular reciprocal inhibition is invoked to explain the observation that vestibular stimulation can disrupt performance on visualisation and mental rotation tasks. Note that although vision is the critical sensory modality for the normal calibration of vestibular signals, the vestibular system can utilise non-visual sensory signals as evidenced by the ability of congenitally blind individuals to orientate themselves using only vestibular cues of motion.

The neural basis for the higher order (perceptual) brain response during visuo-vestibular interaction has been explored using functional imaging, lesion mapping and brain stimulation. Unlike the motor or somatosensory systems, there is no primary vestibular cortex, rather vestibular signals are widely conveyed to the cerebral cortex. Conversely vestibular sensitive cortical neurones invariably display reactivity to other sensory inputs such as proprioception or visual motion, i.e. vestibular sensitive neurones are truly multi-modal sensory neurones.

Vestibular stimulation e.g. via bithalamic caloric irrigation or galvanic stimulation of the vestibular nerve, is associated with increased neuro-imaging signal in a network of brain regions primarily in the Sylvian fissure, insula, retroinsular cortex, fronto-parietal operculum, superior temporal gyrus and cingulate cortex. Conversely, signal reduction is observed in visual cortex (the neuro-imaging correlate of visual-vestibular reciprocal inhibition). In contrast optokinetic visual stimulation inducingvection engenders an opposite pattern, i.e. reduced signal in somatosensory and parietal (‘vestibular’) areas versus increased signal in visual cortex.

Since neuroimaging is a correlational technique we recently utilised transcranial magnetic stimulation (TMS) to probe visual cortical excitability during vestibular activation (see Figure 1B and reference 12). We posed two main questions: first, is there a true change in visual cortical excitability during vestibular activation; second, is the visual cortical response uniform or is there a differential response between early visual cortex (includes V1/2) and visual motion cortex (area V5/MT). This latter aspect was prompted by the lack of clarity in the literature with some suggesting a uniform visual cortical involvement, versus those proposing a selective involvement of visual motion areas. In this experiment, we probed the excitability of visual motion area V5/MT and separately early visual cortex (EVC), i.e. areas V1 and V2, using TMS, during vestibular activation (obtained by caloric stimulation). TMS can be used to probe visual cortical excitability by measuring the relative ease with which one can evoke a phosphene (a perceived flash of light elicited by visual cortex electrical or magnetic stimulation). We found that vestibular stimulation was associated with decreased V5/MT excitability versus increased excitability of early visual cortex (see Figure 1B). Thus, strong stimulation of the vestibular system may reduce sensitivity of visual motion detection areas, but crucially leaves early visual cortex functionally intact (thus not interfering with visual discriminative functioning). This finding provides a possible neurophysiological correlate for the putative reciprocal inhibition between vestibular and visual cortical networks.

Functional imaging has also been used to compare the brain activation of healthy controls during vestibular stimulation to activity in patients following a vestibular lesion. Patients with vestibular neuritis were examined using positron-emission tomography during the acute stage and again three months later.

Figure 1. A) The ‘train effect’. To differentiate between self versus object motion, visual information alone is sometimes insufficient, therefore the brain can also employ information from the vestibular system to provide an estimate of absolute motion of the head. B) During caloric stimulation the probability of perceiving a phosphene (a marker of visual cortical excitability) was significantly reduced in V5/MT regions, whereas the opposite effect was observed in the EVC (see reference 12). C) The rotating disc (left) affects the perception of the ‘true’ gravitational vertical in the direction of tilt. The extent to which an individual’s perception is influenced by these backgrounds provides a measure of visual dependence. The planetarium (right) is used as a rehabilitation therapy for patients with visual vertigo (modified from reference 51).
Increases in regional cerebral glucose metabolism (rCGM) were found in the acute stage relative to the recovery period in multisensory vestibular cortical areas, whereas reduced rCGM was reported in visual and somatosensory areas. These acute stage activation patterns largely mirror those described during vestibular stimulation in healthy volunteers. The relationship between brain structure and vestibular function has also been investigated using neuroimaging. In a follow-up study, patients with VN were tested at least six months after disease onset. Increased grey matter density was reported in medial vestibular cortical areas, whereas reduced grey matter was found in left hippocampus and right superior temporal gyrus. Patients who reported residual canal paresis also showed increased grey matter density bilaterally in visual-motion sensitive areas in middle temporal cortex (MT/V5), an area that also receives vestibular input. This may reflect an attempt to compensate for (vestibular) motion sensitive deficits experienced by patients with significant vestibular deficits after vestibular neuritis. Outcomes as measured by the clinical vestibular score and subjective vestibular disability score were positively correlated with grey matter density in insular, retroinsular and MT and STG regions. These studies indicate that both brain functional and structural changes may take place during central vestibular compensation. Similarly, functional and imaging changes in visual mechanisms develop in patients with bilateral vestibular failure (e.g. secondary to gentamicin or idiopathic), probably underpinning adaptation to the oscillopsia experienced by bilateral vestibular patients.

Clinical Relevance – ‘Visually-induced dizziness’

The visuo-vestibular interaction is of particular clinical relevance to patients suffering from visually-induced dizziness, previously known as visual vertigo, ‘visuo-vestibular mismatch’ or ‘space and motion discomfort’. Patients with visually-induced dizziness report dizziness, unsteadiness and disorientation in visually disorienting surroundings but typically not classical rotational vertigo. The distinguishing characteristic of these patients compared to other dizziness patients is their tendency to be over reliant upon vision for postural control and balance, a situation termed ‘visual dependency’. Visually-induced dizziness appears to be the end result of repeated exposure to dizziness developing in diagnoses as disparate as vestibular migraine (see consensus statement on diagnosis of vestibular migraine) BPPV or post-vestibular neuritis. Visually-induced dizziness occurring post-vestibular neuritis should be distinguished from symptoms related to ‘poor compensation’ of the acquired unilateral peripheral vestibular deficit due to the vestibular neuritis. Poor compensation from a post-vestibular neuritis vestibular deficit can arise due to intermittent vertigo from BPPV, physical inactivity and/or excessive vestibular sedative therapy. A clear identification of the triggers (visual and non-visual) of symptoms post-vestibular neuritis is important since this determines the therapeutic intervention to alleviate these symptoms, e.g. stopping vestibular sedatives, initiating anti-migraine drugs, positional manoeuvres for BPPV, and vestibular physiotherapy. In a patient with post-vestibular neuritis visually-induced dizziness, typically as the patient’s overt continuous spinning vertigo abates another form of dizziness gradually increases whereby dizziness occurs in visually-busy situations such as supermarkets or shopping malls (leading to the false conclusion that the patient has a primary agoraphobia). Sometimes the patient does not discriminate between the acute attack of vertigo and subsequent chronic dizziness, and reports an acute onset with failure to improve over subsequent months. These patients may fail to report visual motion exacerbation of symptoms so it is important to always ask about ‘supermarkets’, ‘shopping malls’, ‘moving trains’, ‘action films’ or ‘video games’ as triggers of dizziness.

In Figure 2 we outline a hypothetical schema of the brain mechanisms underlying visual-vestibular interaction and visually-induced dizziness. An acute vestibular lesion results in an unreliable vestibular signal leading to impaired visual and postural stability. This unreliable vestibular signal is partially corrected for by the brain shifting its reliance from vestibular to visual motion infor...
mation (‘visual dependency’). This acute visual
dependence usually remits once the tonic
vestibular imbalance resolves (via a process of
rapid brainstem plasticity). Occasionally, this
visual dependence persists despite an
adequate rebalancing of the vestibular signal
leading to a maladaptive state of visually-
induced dizziness. Why some patients go on to
develop long term visual dependence and
vestibularly induced dizziness is not completely
understood however investigation of the mecha-

nisms of brain plasticity responsible for these
symptoms are on-going.46

Whatever the neurobiological mechanisms
underlying visually-induced dizziness and visual
dependency important aggravators of
vestibularly induced dizziness include psychologi-
cal symptoms,43 and migraine. How these
stimuli is an effective treatment.47,48 The first
evidence from experiments in both healthy
vestibular patients, presumably by impairing
sedatives is inimical to the recovery of
chronic treatment (>3 days) with vestibular
the adaptive change required for symptomatic
improvements in visual vertigo symptom
rehabilitation. Interventions included a plane-
tary and optokinetic disc stimuli (See
Figure 2C) in order to examine whether visual
training and exposure to visual stimuli may
improve the symptoms of chronically dizzy
patients by addressing the imbalance in their
visuo-vestibular interaction and visual depend-
cy. It is recommended that dizzy patients also
pursue behaviours which challenge their visual
dependency in addition to any formal rehabili-
tation they take part in. This is important as ‘real-
world’ phenomena can never be fully repli-
cated in the lab. Particularly helpful sports
include those requiring VOR-smooth pursuit
integration, e.g. ball sports such as tennis.

Clinical overview
An understanding of visuo-vestibular interac-
tion and the underlying brain mechanisms is
key in understanding patients’ superficially
bizarre complaints (‘I feel dizzy when faced with
shopping mall crowds or walking down
supermarket aisles’) and secondly in devel-
oping effective treatment for visual vertigo.
One potentially problematic group are
vestibular migraineurs who frequently also
complain of visually-induced dizziness. When
treating such patients it is imperative to follow
a step-wise approach. The first step is to treat
the migraine with effective prophylaxis. We
find that standard anti-migrainous drugs work
well with propanolol being our first line
(second line according to patient profile;
including amitriptyline, topiramate, sodium
valporate or pizotifen). Often simply treating
the vestibular migraine with pharmacotherapy
improves the visual symptoms as well. If visually-
induced dizziness persists despite good
migraine control, we then initiate OKN therapy. If
OKN is provided to active migraineurs then
symptoms can be aggravated, hence the
importance of the first step (in treating the
migraine). Once commenced on effective anti-
migraine prophylaxis OKN therapy can be
provided if symptoms of of visually-induced
dizziness persist. Indeed migraineurs show the
greatest improvement in response to OKN
therapy compared to patients with other
chronic peripheral vestibular symptoms.46
Note however that the clinician should be
alert to patients with psychological symptoms
who also avoid visually busy environments for
different reasons, e.g. agoraphobia (ref50).
Equally many vestibular patients suffer from
psychological symptoms as a result of their
vestibular symptoms. In cases of doubt a
liaison psychiatric opinion should be sought.
Needless to say, some patients require a two
pronged vestibular and psychological therapy
approach. As always in medicine, a diagnosis
and appropriate treatment has to be decided
on multiple aspects of the clinical history and
investigations.

Conclusion
An understanding of the brain mechanisms
mediating visual and vestibular interaction has
been little studied however multi-modal
research involving neuroimaging lesion
mapping and more recently TMS has enabled a
mechanistic explanation for patients’ symptoms and the logical develop-
ment of their treatment. There are many unan-
swered pertaining to the modulators of visual-
vestibular interaction, such as migraine,
anxiety and co-existing medical and neurolog-
ical disorders.

REFERENCES
2. Warspe W & Henn V. Neural activity in the vestibular
nucleus of the alert monkey during vestibular and optoki-
4. Palla A, Straumann D. & Bronstein AM. Vestibular
neuritis: vertigo and the high-acceleration vestibular-ocular
5. Del Orio LF & Leigh RJ. Fixation period stability and
oculopulse suppression in congenital nystagmus: an
6. Poldst J, Brandt T & Degos D. Objective motion detection
7. Poldst J, Straube A & Bings W. Differential effects of
ambisensory vestibulo-sensitomotor stimulation on the
8. Mast PW, Merfeld DM & Koslow SM. Visual mental
imagery during caloric vestibular stimulation. Neuropsychologia
Vestibular perception and navigation in the congenitally
11. Kadi J, Bronstein AM, Mahabat P, Nigmatullina Y &
Seemungal BM. Humans use an internal clock for esti-
imating their position in space. In Society for
Neuro-otological emergencies.

12. Seemungal BM, Guzman-Lopez J, Arndt G, Schultz SR,
Walsh V & Yusof N. Vestibular Activation Differentially
Modulates Human Early Visual Cortex and V5/MT.
Neural correlates of visual-motion perception as object or self-motion. Neurosciology
2012; 16:873-82.
13. Grüsser D, Paus M & Schneider G. Localization
and responses in the posterior-inferior vestibular
cortex of awake monkeys (Macaca fuscata).
The journal of physiology 1990;430:537-57.
14. Guldin WO & Grüsser OJ. Is there a vestibular cortex?
15. Grüsser D, Paus M & Schneider G. Vestibular neurons
in the posterior-inferior vestibular cortex of awake monkeys (Macaca fuscata).
during involuntary ocular oscillations: Brain
New Journal Reviews Editor for ACNR

We would like to welcome Gemma Cummins as our new Journal Reviews editor. Gemma is a Specialist Registrar in Neurology at Addenbrooke’s Hospital, Cambridge and is currently completing a PhD on movement disorders and cognition at the Van Geest Centre for Brain Repair, Cambridge.

Prof Ebers awarded AAN Prize for MS research

The American Academy of Neurology and the National Multiple Sclerosis Society awarded the 2013 John Dystel Prize for MS Research to George C. Ebers, MD, a researcher with the University of Oxford and Oxford University Hospitals Trust in Oxford, UK. Ebers received the award at the Academy’s 65th Annual Meeting in San Diego, earlier this year. The John Dystel Prize recognises a significant contribution to research in the understanding, treatment or prevention of multiple sclerosis (MS). Ebers’ research has focused on genetic and environmental influences on MS risks. “We have found that MS risk factors previously considered to be genetic can be changed based on environment, strongly implicating gene-environment interaction. Our studies highlight how climate and diet relate to factors leading to MS, which can be viewed as a largely preventable disease. Vitamin D exposure appears to be the main factor determining geographical risk” said Ebers.

MND Association Lectureship in Translational Neuroscience

Dr Richard Mead, based at the Sheffield Institute for Translational Neuroscience (SiTraN) at the University of Sheffield, has been awarded the Kenneth Snowman-MND Association Lectureship in Translational Neuroscience.

The five-year Kenneth Snowman-MND Association lectureship is aimed to embed pre-clinical expertise in motor neuron disease (MND) models within SiTraN as a national resource.

Dr Richard Mead was awarded the lectureship as he has the expertise and knowledge to enable high quality pre-clinical research into MND.

Dr Mead has over 14 years experience in both academia and industry with a background in models of MND (mice and fibroblasts or ‘skin cells’) and multiple sclerosis.

Developing disease models is important for furthering our understanding of MND and allows researchers to screen potential new drugs for a beneficial effect before they can be given to humans, by means of a clinical trial.

As well as a track record of taking compounds into clinical development, Dr Mead hopes to use this knowledge and experience to develop MND specific therapeutic compounds. Dr Mead has already shown effectiveness of two compounds using his pre-clinical screening programme, with one being given ‘Orphan drug’ designation by the European Medicines Agency (EMA).

For more information see www.mndassociation.org

WFNR Francis Gerstenbrand Award deadline

1st November 2013 is the deadline for entries for the World Federation for Neurorehabilitation (WFNR) Francis Gerstenbrand Award. The Award is open to clinicians, researchers and allied health professionals and recognises and rewards a neurorehabilitation project that has benefitted patients. The annual, single prize of £3000 will be awarded as either a travel bursary to a clinical conference, professional development course or research project.

Named after Professor Francis Gerstenbrand, in recognition of his continuous contributions to neurorehabilitation, the Award is open to WFNR members and non-members worldwide.

Entries can come from any aspect of neurorehabilitation and examples include a patient or clinic management initiative, research project, best practice development or the use of a new technological development. A panel of four or five judges, led by the WFNR President, will review the entries.

For further details and details on how to apply, visit: http://wfnr.co.uk/en/education-and-research/wfner-award/